Amendments to the Specification:

Please replace paragraph number [0096] with the following rewritten paragraph:

These results indicate that many neural precursor lineages respond similarly to the overexpression of c-myc. In addition to primary neural cultures prepared from nervous system tissues
of mammals, recent advances in embryonic stem cell cultures indicate that various neural
precursors form in vitro during differentiation of totipotential or pluripotent embryonic stem cells
and cell lines maintained in culture for long term (Renoncourt et al., Mech. Dev. (1998) 78, 185;
Svendsen et. al., Trends Neurosci. (1999) 22, 357; Brustle et. al., Science (1999) 285, 754.).
These cultures can generate nestingnestin-positive neural precursor cells which can then be
transferred to serum-free medium and subsequently expanded with bFGF and/or EGF for short
term. Long-term, mass expansion has not been feasible since the initial neural precursor
formation is inefficient. However, by utilizing the genetic modification method with c-myc gene
described here, those transient neural precursors may be turned into stable cell lines.